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Coccidiosis poultry vaccine

Field of the Invention

The present invention relates to a protein derived from Eimeria acervulina, which is capable of stimulating immune lymphocytes. It also relates to a nucleic acid sequence encoding all or an antigenically significant part of this protein, a recombinant vector comprising such a nucleic acid sequence, a host cell or organism transformed with such a recombinant vector and a vaccine for the protection of poultry against coccidiosis.

A Background of the Invention

Coccidiosis is a disease caused by infection with many species of coccidia, of the more one or intracellular protozoal parasites of the Apicomplexa and the genus Eimeria. Poultry is defined herein as domesticated birds that serve as a source of include such commercially meat and that eggs or important kinds as chickens, turkeys, ducks, geese, quinea fowl, pheasants, pigeons and peafowl.

Coccidiosis in chickens is known to be caused by several different species of Eimeria, namely Eimeria acervulina, E. maxima, E. tenella, E. necatrix, E. brunetti, E. mitis, E. praecox, E. mivati and E. hagani. Some people, however, doubt the true existence of the last two species. Low level infection with any of these Eimeria species results in a protective immunity to reinfection.

The species do differ in their pathogenic effect on chickens, the type of chicken also playing a role; thus, a broiler chicken will be subjected to a great deal of damage by a parasite such as E. acervulina or E. maxima because these parasitise large portions of

the small intestine, where food digestion plays a major role.

E. acervulina is one of the most common species found in the litter of broiler houses in both Europe and the USA. It has a great reproductive potential and is regarded as pathogenic because it produces a marked depression in gain of body weight, higher feed conversion and it produces gross lesions in the upper small intestine.

During the life cycle (see also Table 1), Eimeria parasite passes through a number of stages. The life cycle begins when the chicken ingests the known as the sporulated oocyst, infectious stage, during ground feeding or by inhalation of dust. of the sporulated oocyst is ruptured by a combination of mechanical grinding action and chemical action in the gizzard and intestinal tract, resulting in the release of four sporocysts. The sporocysts pass into the duodenum where they are exposed to bile and digestive enzymes resulting in the release of two sporozoites per sporocyst.

Table 1. Endogenous stages of Eimeria acervulina in stained sections of infected duodenum (after McDonald V. et al., Parasitol. 8, 21-30, 1982).

Time of infection	Histological observations
24 h	Immature 1 St generation asexual stages
36 h	Semi-mature 1 St generation schizonts
42 h	Mature 1 st gen. schizonts. Immature 2 nd gen. parasites
48 h	Mature 2 nd gen. schizonts. A few 3 rd gen. schizonts with 8-16 merozoites
60 h	Mature 3 rd gen. schizonts, immature 4 th gen. parasites

The sporozoites are mobile and search for suitable host epithelium cells in order to penetrate and reproduce in them. Following infection of an

epithelium cell, the parasite enters the schizont phase of its life cycle, producing from 8 to 16 to >200 merozoites per schizont. Once released from the schizont, the merozoites are free to infect further epithelium cells. After from two to five of these intracellular the reproduction cycles, asexual merozoites grow into sexual forms known as the female or macrogametocyte and the male or microgametocyte. Following fertilization of the macrogametocyte by the microgametes released from the microgametocyte, zygote is formed which creates a cyst wall about itself. The newly formed oocyst is passed out of the infected chicken with the droppings.

conditions of environmental correct With the temperature and humidity and sufficient oxygen in the air, the oocyst will sporulate into the infectious infect new host and ready to a spreading the disease. Thus no intermediate host is required for transfer of the parasite from bird to bird.

The result of the Eimeria parasite infecting the digestive tract of a chicken may be a reduction in weight gain, increased feed conversion, cessation of egg production and, in some cases, death. The increase intensive production of poultry in accompanied by severe losses due to this parasite; indeed, coccidiosis has become the most economically important parasitic disease. In the Netherlands, the losses that poultry farmers suffer every year run into millions of guilders; in 1986 the loss was about 13 million guilders. In the same year, a loss of 300 million dollars was suffered in the United States.

In the past, several methods have been used in attempts to control coccidiosis. Prior to the advent

of chemotherapeutic agents, improved sanitation using disinfectants, together with the mechanical removal of litter, was the main method employed; sufficient oocysts, however, usually remained to transmit the disease.

The introduction of coccidiostatic agents in the drinking water, in addition some success at disease in management, resulted control. Such agents have been found to suffer from a drop in effectiveness over the years, due partly to the development of drug resistant strains of coccidia. Furthermore, several chemotherapeutic agents have been leave residues in the meat, making it found to unsuitable for consumption.

Attempts have been made to control the disease immunologically by administering to chickens a live vaccine comprising oocysts from all seven species of Eimeria. the oocysts administered being precocious lines. Such precocious lines are obtained by inoculating chickens with a wild population of an collecting the very Eimeria species and are excreted as a result parasites that infection. The collected parasites are put back into chickens and the cycle is repeated several times. Eventually a precocious line of parasite is produced which has fewer cycles of asexual reproduction in the Thus such lines retain their immunogenicity, whilst producing fewer parasites in the gut with less consequential damage being caused to the host chicken.

The disadvantage of this type of vaccine is that it is expensive to produce because of the necessity of producing it in live chickens and its lower reproductive potential.

The advent of genetic engineering has provided new methods for producing effective vaccines. Using these methods, the DNA coding for the antigenic proteins of some pathogenic microorganisms has been cloned into Escherichia coli microorganisms as host such Salmonella spec., with the result that the protein has been expressed at sufficiently high levels such that it can be incorporated into a vaccine. The advantage of proteins produced in this way is that they are noninfectious and are relatively cheap to produce. In this way, vaccines have been prepared against a number of viruses such as hepatitis, herpes simplex and foot and mouth disease.

Attempts have been made to genetically engineer a coccidiosis vaccine. European patent application No. 337 589 describes the isolation of a Group B Eimeria its insertion into tenella protein and expression vector which, in turn, has been used to transform appropriate hosts. Patent Cooperation Treaty Application WO 92/04461 describes the construction of a microorganism that produces an antigenic protein using either the "mRNA route" or the "nuclear route". In this way, certain antigens from E. tenella and E. maxima were prepared and sequenced. Taking this type of route to prepare antigens for incorporation into a vaccine relies only upon selecting antigens induce antibodies in heterologous which could an species. This approach does not necessarily end up with selecting the most protective antigen.

From H.S. Lillehoj (Vet. Immunol. Immunopath., 13, 321-330, 1986) it can be conceived that development of protective immunity in chickens infected with coccidia may be due to the development of a species-specific T cell response.

A Summary of the Invention

It has now been found that a very immunogenic protein can be isolated from the 42hr developmental stage of Eimeria schizonts. Surprisingly, this protein is found intracellularly in Eimeria and it appears to contain high sequence homology with known heterologous lactate dehydrogenases (LDH).

Thus, the invention provides a protein having one or more immunoreactive and/or antigenic determinants of Eimeria lactate dehydrogenase, which has a monomeric molecular weight of about 37 kD.

More specifically the lactate hydrogenase is derived from Eimeria acervulina.

According to a second aspect of the invention, there is provided a nucleic acid sequence encoding all part, in particular substantial immunologically active part, of a purified Eimeria lactate dehydrogenase. Such a nucleic acid sequence expression be operatively linked to sequences resulting in a recombinant nucleic acid molecule which, when inserted into a suitable vector, results in a recombinant vector capable of expressing the nucleic acid sequence.

Such a recombinant vector, or nucleic acid sequence as defined above, may be used to transform a suitable host cell or organism. Such a transformed host cell or organism may, in turn, be used to produce the stimulatory protein for incorporation into a vaccine

for the protection of poultry against coccidiosis. Alternatively, the transformed host cell or organism may itself be incorporated into a vaccine.

A Detailed Description of the Envention refers "protein" term general, with biological molecular chain amino acids of activity. A protein is not of a specific length and can, if required, be modified in vivo or in vitro, by, amidation, carboxylation for example, glycosylation, thus, inter alia, peptides, phosphorylation;

oligopeptides and polypeptides are included within the

definition.

More particularly, this invention provides proteins possessing LDH activity, or immunogenically active parts thereof, which have the amino acid sequence shown in SEQ ID NO. 2 and their biologically functional equivalents or variants.

The biologically functional equivalents or variants of the proteins specifically disclosed herein are proteins derived from the abovenoted amino acid sequences, for example by deletions, insertions and/or substitutions of one or more amino acids, but retain one or more immunogenic determinants of the Eimeria antigens, i.e. said variants have one or more epitopes capable of eliciting an immune response in a host animal.

It will be understood that, for the particular proteins embraced herein, natural variations can exist between individual Eimeria parasites or strains. These variations may be demonstrated by (an) amino acid difference(s) in the overall sequence or by deletions, substitutions, insertions, inversions or additions of in said sequence. Amino acid amino acid(s) (an) essentially alter do not substitutions which

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immunological activities, have been biological and described, e.g. by Neurath et al in "The Proteins" acid (1979).Amino York New Academic Press acids or amino related between replacements frequently have occurred replacements which evolution are, inter alia, Ser/Ala, Ser/Gly, Asp/Gly, Asp/Asn, Ile/Val (see Dayhof, M.D., Atlas of protein sequence and structure, Nat. Res. Found., Biomed. Washington D.C., 1978, vol. 5, suppl. 3). Other amino acid substitutions include Asp/Glu, Thr/Ser, Ala/Gly, Ala/Thr, Ser/Asn, Ala/Val, Thr/Phe, Ala/Pro, Lys/Arg, Based and Ala/Glu. Leu/Val Leu/Ile, information, Lipman and Pearson developed a method for rapid and sensitive protein comparison (Science, 227, functional determining the and 1435-1441, 1985) similarity between homologous proteins. Such acid substitutions of the exemplary embodiments of this invention are within the scope of the invention resulting proteins retain as the long immunoreactivity.

immunogenic fragments the also Furthermore, herein orspecifically disclosed proteins present in the included functional variants are invention.

The term "fragment" as used herein means a DNA or amino acid sequence comprising a subsequence of the nucleic acid sequence or protein of the invention. Said fragment is or encodes a polypeptide having one or more immunogenic determinants of an Eimeria antigen. Methods for determining usable immunogenic polypeptide fragments are outlined below. Fragments can inter alia be produced by enzymatic cleavage of precursor molecules, using restriction endonucleases for the DNA and proteases for the polypeptides. Other methods include chemical synthesis of the fragments or

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the expression of polypeptide fragments by DNA fragments.

Suitable immunogenic polypeptide fragments protein according to the invention containing (an) by means of the method be found epitope(s) can described in Patent Application WO 86/06487, Geysen, 3998-4002, 81, et al. (Proc. Natl. Acad. Sci. Immunol. Meth. 1984), Geysen, H.M. et al. (J. 259-274, 1987) based on the so-called pepscan method, wherein a series of partially overlapping peptides corresponding with partial sequences of the complete polypeptide under consideration, are synthesized and their reactivity with antibodies is investigated.

addition, a number of regions of the polypeptide, with the stated amino acid sequence, can be theoretical of the basis epitopes on designated considerations and structural agreement with epitopes The determination of now known. which are of the combination is based on a regions hydrophilicity criteria according to Hopp and Woods (Proc. Natl. Acad. Sci. 78, 3824-3828, 1981) and the secondary structure aspects according to Chou Fasman (Advances in Enzymology 47, 45-148, 1987).

T-cell epitopes which may be necessary can likewise be derived on theoretical grounds, e.g. with the aid of Berzofsky's amphiphilicity criterion (Science 235, 1059-62, 1987).

and invention further provides isolated The purified nucleic acid sequences encoding the above mentioned proteins of Eimeria. One of these nucleic acid sequences is shown in SEQ. ID. NO. 1. It is well known in the art that the degeneracy of the genetic the of bases in substitution permits resulting in another codon but still coding for the same amino acid, e.g. the codon for the amino acid glutamic acid is both GAT and GAA. Consequently, it is

clear that, for the expression of a protein with the amino acid sequence shown in SEQ. ID. NO. 2, the nucleic acid sequence may have a codon composition different from the nucleic acid sequence shown in SEQ. ID. NO. 1.

A nucleic acid sequence according to the present invention may be isolated from an Eimeria strain and multiplied by recombinant DNA techniques including polymerase chain reaction (PCR) technology or may be chemically synthesized in vitro by techniques known in the art.

A nucleic acid sequence according to the invention can be ligated to various replication effecting DNA sequences with which it is not associated, or linked in nature, resulting in a so-called recombinant vector which can be used for the transformation of a suitable host. Useful recombinant vectors are preferably derived from plasmids, bacteriophages, cosmids or viruses.

Specific vectors or cloning vehicles which can be used to clone nucleic acid sequences according to the invention are known in the art and include inter alia plasmid vectors such as pBR322, the various pUC, pGEM and Bluescript plasmids; bacteriophages, e.g. \alphagtat-Wes, Charon 28 and the M13 derived phages or viral vectors such as SV40, adenovirus or polyoma virus (see also Rodriguez, R.L. and D.T. Denhardt, ed., Vectors: A survey of molecular cloning vectors and their uses, J.A. al., Butterworths, 1988; Lenstra, et Virol., 110, 1-24, 1990). The methods to be used for the construction of a recombinant vector according to the invention are known to those of ordinary skill in the art and are inter alia set forth in Maniatis, T.

et al. (Molecular Cloning A Laboratory Manual, second edition; Cold Spring Harbor Laboratory, 1989).

For example, the insertion of the nucleic acid sequence according to the invention into a cloning vector can easily be achieved when both the genes and the desired cloning vehicle have been cut with the same restriction enzyme(s) as complementary DNA termini are thereby produced.

Alternatively, it may be necessary to modify the restriction sites that are produced into blunt ends either by digesting the single-stranded DNA or by filling in the single-stranded termini with an appropriate DNA polymerase. Subsequently, blunt end ligation with an enzyme such as T4 DNA ligase may be carried out.

If desired, any restriction site may be produced by ligating linkers onto the DNA termini. Such linkers may comprise specific oligonucleotide sequences that encode restriction site sequences. The restriction enzyme cleaved vector and nucleic acid sequence may also be modified by homopolymeric tailing.

"Transformation", as used herein, refers to introduction of an heterologous nucleic acid sequence into a host cell, irrespective of the method used, for transduction. uptake orexample direct heterologous nucleic acid sequence may be maintained through autonomous replication or, alternatively, may be integrated into the host genome. If desired, the recombinant vectors are provided with appropriate control sequences compatible with the designated host. These sequences can regulate the expression of the addition to inserted nucleic acid sequence. In

microorganisms, cell cultures derived from multicellular organisms may also be used as hosts.

The recombinant vectors according to the invention preferably contain one or more marker activities that may be used to select for desired transformants, such as ampicillin and tetracycline resistance in pBR322, ampicillin resistance and α -peptide of β -galactosidase in pUC8.

A suitable host cell is a microorganism or cell which can be transformed by a nucleic acid sequence encoding a polypeptide or by a recombinant vector comprising such a nucleic acid sequence, and which can, if desired, be used to express said polypeptide encoded by said nucleic acid sequence. The host cell can be of prokaryotic origin, e.g. bacteria such as Escherichia coli, Bacillus subtilis and Pseudomonas species; or of eukaryotic origin such as yeasts, e.g. Saccharomyces cerevisiae or higher eukaryotic cells such as insect, plant or mammalian cells, including HeLa cells and Chinese hamster ovary (CHO) Insect cells include the Sf9 cell line of Spodoptera frugiperda (Luckow et al., Biotechnology 1988). Information with respect to the cloning and expression of the nucleic acid sequence of the present invention in eukaryotic cloning systems can be found in Esser, K. et al. (Plasmids of Eukaryotes, Springer-Verlag, 1986).

In general, prokaryotes are preferred for the construction of the recombinant vectors useful in the present invention. E.coli K12 strains are particularly useful, especially DH5a or MC1061 strains.

For expression, nucleic acid sequences of the present invention are introduced into an expression

vector, i.e. said sequences are operably linked to expression control sequences. Such control sequences operators, promotors, enhancers, comprise inducers, ribosome binding sites etc. Therefore, the recombinant provides a invention present comprising a nucleic acid sequence encoding an Eimeria protein identified above operably linked to expression control sequences, which is capable of expressing the DNA sequences contained therein in (a) transformed host cell(s).

It should be understood, of course, that the nucleotide sequences inserted at the selected site of the cloning vector may include nucleotides which are not part of the actual structural gene for the desired polypeptide, or may include only a fragment of the complete structural gene for the desired protein as long as the transformed host will produce a polypeptide having at least one or more immunogenic determinants of an Eimeria protein antigen.

When the host cells are bacteria, useful expression control sequences which may be used include the Trp promotor and operator (Goeddel, et al., Nucl. Acids Res., 8, 4057, 1980); the lac promotor and operator (Chang, et al., Nature, 275, 615, 1978); the outer membrane protein promotor (Nakamura, K. and Inouge, M., EMBO J., 1, 771-775, 1982); the bacteriophage lambda promotors and operators (Remaut, E. et al., Nucl. Acids Res., 11, 4677-4688, 1983); the α -amylase subtilis) promotor and operator, termination (B. sequences and other expression enhancement and control sequences compatible with the selected host cell. When the host cell is yeast, illustrative useful expression control sequences include, e.g., α -mating factor. For insect cells the polyhedrin or p10 promotors baculoviruses can be used (Smith, G.E. et al., Mol.

Cell. Biol. 3, 2156-65, 1983). When the host cell is mammalian origin illustrative useful expression control sequences include the SV-40 promotor (Berman, Science, 222, 524-527, 1983) P.W. et al., metallothionein promotor (Brinster, R.L., Nature, 296, 39-42, 1982) or a heat shock promotor (Voellmy et al., 4949-53, 1985). Sci. USA, 82, Natl. Acad. Alternatively, expression control sequences present in For maximizing Eimeria may also be applied. expression, see also Roberts and Lauer (Methods in Enzymology, 68, 473, 1979).

Therefore, the invention also comprises (a) host cell(s) containing a nucleic acid sequence or a recombinant nucleic acid molecule or a recombinant vector described above, capable of producing the Eimeria protein by expression of the nucleic acid sequence.

Immunization of poultry against Eimeria infection administering to the birds can be achieved by in invention an according to the protein immunologically relevant so-called context as a subunit vaccine. The subunit vaccine according to the invention may comprise a protein in a pure form, a pharmaceutically in the presence of optionally The protein can optionally acceptable carrier. covalently bonded to a non-related protein, which can be of advantage in the purification of the fusion Examples are B-galactosidase, protein product. prochymosine, blood clotting factor Xa, etc.

In some cases the ability to raise protective immunity using these proteins per se may be low. Small fragments are preferably conjugated to carrier molecules in order to raise their immunogenicity.

Suitable carriers for this purpose are macromolecules, polymers (proteins like key such as natural synthetic toxins), hemocyanin, albumin, limpet (polylysine, acids polyamino like polymers amphiphilic compounds micelles of polyalanine), or like saponins. Alternatively these fragments may be linear thereof, preferably as polymers provided polymers.

proteins according the the Ιf required, invention which are to be used in a vaccine can be or in vivo, for by vitro in glycosylation, acylation, amidation, carboxylation or phosphorylation.

A newly developed vaccine version is a vaccine in which the DNA coding for the protein of the invention is administered in a pharmaceutically acceptable form, for instance in the form of "bullets", which can be shot into the tissue. This naked DNA can be used as vaccine provided it is presented in a plasmid or in eukaryotic promoter suitable with combination sequences such as those from SV40 virus. In this way one can achieve the introduction of this DNA into the thus ensuring the expression of genomic DNA, antigen in situ.

is live subunit vaccines to alternative vaccines. A nucleic acid sequence according to invention is introduced by recombinant DNA techniques into a microorganism (e.g. a bacterium or virus) such a way that the recombinant microorganism is still able to replicate, thereby expressing a polypeptide acid sequence inserted nucleic by the eliciting an immune response in the infected host bird.

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A preferred embodiment of the present invention is a recombinant vector virus comprising an heterologous nucleic acid sequence described above, capable of expressing the DNA sequence in (a) host cell(s) or host bird infected with the recombinant vector virus. The term "heterologous" indicates that the nucleic acid sequence according to the invention is not normally present in nature in the vector virus.

Furthermore, the invention also comprises (a) host cell(s) or cell culture infected with the recombinant vector virus, capable of producing the Eimeria protein by expression of the nucleic acid sequence.

For example the well known technique of in vivo homologous recombination can be used to introduce an heterologous nucleic acid sequence according to the invention into the genome of the vector virus.

DNA fragment corresponding First, a insertion region of the vector genome, i.e. a region used for the incorporation which can be sequence without disrupting heterologous functions of the vector such as those necessary for infection or replication, is inserted into a cloning according to standard recDNA techniques. vector Insertion-regions have been reported for number of microorganisms (e.g. EP 80,806, EP 110,385, EP 83,286, EP 314,569, WO 88/02022, WO 88/07088, US 4,769,330 and US 4,722,848).

Second, if desired, a deletion can be introduced into the insertion region present in the recombinant vector molecule obtained from the first step. This can be achieved for example by appropriate exonuclease III digestion or restriction enzyme treatment of the recombinant vector molecule from the first step.

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the heterologous nucleic acid sequence insertion region present into the recombinant vector of the first step or in place of from said recombinant vector. The deleted insertion region DNA sequence should be of appropriate length as to allow homologous recombination with the vector genome to occur. Thereafter, suitable cells can be infected with wild-type vector virus or transformed of the genomic DNA in the presence vector recombinant vector containing the insertion flanked by appropriate vector DNA sequences whereby recombination corresponding occurs between the regions recombinant vector and the vector genome. Recombinant vector progeny can now be produced in cell culture and example genotypically be selected for can by hybridization, detecting phenotypically, e.g. enzyme activity encoded by a gene co-integrated along acid heterologous nucleic sequence, the polypeptide antigenic heterologous detecting the expressed by the recombinant vector immunologically.

microorganism A recombinant microorganisms be this administered to poultry for immunization whereafter it maintains itself for some time, or even replicates in the body of the inoculated animal, expressing in vivo a polypeptide coded for by the inserted nucleic acid sequence according to the invention resulting in the stimulation of the immune system of the inoculated animal. Suitable vectors for the incorporation of a nucleic acid sequence according to the invention can be derived from viruses such as pox viruses, 83,286, US 4,769,330 vaccinia virus (EP 110,385, \mathbf{EP} or fowl pox virus (WO 88/02022), and US 4,722 848) such as HVT adenourus adeno virus (WO 88/07088) or Marek's herpes viruses influenza virus, Disease virus, or specific Salmonella coli bacteria such as Ε. or

species. With recombinant microorganisms of this type, the polypeptide synthesized in the host animal can be exposed as a surface antigen. In this context fusion polypeptide with OMP proteins, the proteins of for example E. coli or synthetic provision of signal and anchor sequences which are recognized by the organism are conceivable. It is also possible that the Eimeria polypeptide, if desired as part of larger whole, is released inside the animal to immunized. In all of these cases it is also possible immunogenic products will ormore one that expression which generate protection against various pathogens and/or against various antigens of a given pathogen.

A vector vaccine according to the invention can be prepared by culturing a recombinant bacterium or a host cell infected with a recombinant vector comprising a nucleic acid sequence according to the invention, whereafter recombinant bacteria or vector containing cells and/or recombinant vector viruses grown in the cells can be collected, optionally in a pure form, and formed into a vaccine optionally in a lyophilised form.

Host cells transformed with a recombinant vector according to the invention can also be cultured under conditions which are favourable for the expression of a polypeptide coded by said nucleic acid sequence. Vaccines may be prepared using samples of the crude host cell lysates or host cell culture, embodiment in another more purified although polypeptides according to the invention are formed depending on its intended use. into a vaccine, order to purify the polypeptides produced, host cells transformed with a recombinant vector according to the invention are cultured in an adequate volume and the

polypeptides produced are isolated from such cells, or protein is excreted. medium if the Polypeptides excreted into the medium can be isolated by standard techniques, e.q. purified ultrafiltration, centrifugation, fractionation, ation or Intracellula affinity chromatography, gel filtration polypeptides cellular whereas A chromatography, isolated by first collecting said cells, disrupting the cells, for example by sonication or by other mechanically disruptive means such as French press, followed by separation of the polypeptides from the other intracellular components and forming the polypeptides into a vaccine. Cell disruption could also be achieved by chemical (e.g. EDTA or detergents Triton X114) or enzymatic means, such as lysozyme digestion.

directed against Antibodies or antiserum а according the invention have to polypeptide diagnostic potential use in passive immunotherapy, anti-idiotypic of immunoassays and generation antibodies.

The Eimeria proteins as characterized above can be antibodies, both to produce used polyclonal, monospecific and monoclonal. If polyclonal antibodies are desired, techniques for producing and processing polyclonal sera are known in the art (e.g. Mayer and Walter. eds, Immunochemical Methods in Cell and Molecular Biology, Academic Press, London, 1987). immunogen antibodies to an can be Monospecific affinity purified from polyspecific antisera modification of the method of Hall et al. (Nature, 379-387, 1984). Monospecific antibody, herein, is defined as a single antibody species or multiple antibody species with homogeneous binding characteristics for the relevant antigen. Homogeneous binding, as used herein, refers to the ability of the antibody species to bind to a specific antigen or epitope.

Monoclonal antibodies, reactive against the Eimeria proteins according to the present invention, can be prepared by immunizing inbred mice by techniques known in the art (Köhler and Milstein, Nature, 256, 495-497, Hybridoma cells are selected by growth in an aminopterin thymidine and hypoxanthine, appropriate cell culture medium such as Dulbecco's modified Eagle's medium. Antibody producing hybridomas are cloned, preferably using the soft agar technique of MacPherson, (Soft Agar Techniques, Tissue Culture Methods and Applications, Kruse and Paterson, eds., Discrete colonies 1973). 276, Academic Press, transferred into individual wells of culture plates for cultivation in an appropriate culture medium. Antibody producing cells are identified by screening Immunogen positive with the appropriate immunogen. hybridoma cells are maintained by techniques known in antibodies Specific anti-monoclonal produced by cultivating the hybridomas in vitro or preparing ascites fluid in mice following hybridoma injection by procedures known in the art.

Anti-idiotypic antibodies are immunoglobulins which carry an "internal image" of the antigen of the pathogen against which protection is desired and can be used as an immunogen in a vaccine (Dreesman et al., J. Infect. Disease, 151, 761, 1985). Techniques for raising anti-idiotypic antibodies are known in the art (MacNamara et al., Science, 226, 1325, 1984).

The vaccine according to the invention administered in a conventional active immunization scheme: single or repeated administration in a manner compatible with the dosage formulation, and in such amount as will be prophylactically effective, i.e. the antigen or recombinant immunizing of amount microorganism capable of expressing said antigen that will induce immunity in poultry against challenge by virulent Eimeria parasites. Immunity is defined as the induction of a significant level of protection in a population of chickens after vaccination compared to an unvaccinated group.

a vaccine protection increase in Next to an comprising the polypeptide of the invention will also reduce the number of oocysts shedded by the infected animals. Normally, the shedded oocysts will other animals in the flock. A decrease in the number of oocysts shedded will then also give a decrease in the number of animals which is subsequently infected and also a decrease in the number of oocysts shedded will give rise to a lesser infective load.

Furthermore, even without effect on the parasite itself, a vaccine can decrease the incidence of disease. This is especially so when the symptoms of the disease are caused by products released by the parasite. Vaccines directed against such products alleviate the symptoms without attacking the parasite.

For live viral vector vaccines the dose rate per chicken may range from $10^5 - 10^8$ pfu. A typical subunit vaccine according to the invention comprises 1 μ g - 1 mg of the protein according to the invention. Such vaccines can be administered intradermally, subcutaneously, intramuscularly, intraperitoneally, intravenously, orally or intranasally.

contain may also vaccine Additionally the aqueous medium or a water containing suspension, often mixed with other constituents in order to increase the activity and/or the shelf life. These constituents may be salts, pH buffers, stabilizers (such as skimmed milk or casein hydrolysate), emulsifiers, adjuvants to oils, response (e.g. immune improve the dipeptide, aluminium hydroxide, saponin, polyanions and amphipatic substances) and preservatives.

A vaccine comprising the polypeptide of the invention may also comprise other immunogenic proteins of E. maxima or immunogenic proteins of other Eimeria species. Such a combination vaccine will decrease the parasitic load in a flock of poultry and will increase the level of protection against coccidiosis.

that а vaccine according clear invention may also contain immunogens related to other pathogens of poultry, or may contain nucleic acid sequences encoding these immunogens, like antigens of Marek's Disease virus (MDV), Newcastle Disease virus etroury Bronchitis virus Infectious Reo virus, Avian Retro virus, Fowl Anemia Agent (CAA),

Adeno virus, Turkey Turkey Rhinotracheitis virus or E. to produce a multivalent vaccine.

The invention is illustrated by the following examples:

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EXAMPLE 1

Handling of parasites

Eimeria acervulina (Houghton strain) and Eimeria tenella (Weybridge strain) parasites were collected after deliberate infection of chickens reared in the absence of coccidia. E. acervulina oocysts were isolated from fecal material on days 4 and 5 post-infection (p.i.). E. tenella oocysts were harvested from the ceca on day 7 p.i..

The oocysts were sporulated with strong aeration at 30°C for 7 hours, resulting in partially sporulated oocysts. Release of sporocysts and sporozoites of 48 hr sporulated oocysts was performed as described earlier in A.N. Vermeulen et al. FEMS Microbiological Letters 110, (1993), 223-230.

intra-cellular stages, To obtain E. acervulina chickens were infected at 5 weeks with 108 sporulated acervulina oocysts. Intracellular parasites were harvested from the duodenum after 42 hours. Hereto chickens were exsanguinated 42hr post inoculation and duodenum was removed from the stomach to Meckel's diverticulum. The tissue was washed and cut into small cm^3 . approximately 1 The pieces pieces of calcium/magnesium free Hanks **BSS** suspended in containing 10 mg/ml glucose (CMF-Hanks). **Epithelial** from the matrix by _ 10 cells were released incubation in EDTA (2 mM EDTA in CMF Hanks at 37°C). Supernatants of four incubations were pooled and centrifuged 10 min at 750g, which pelleted the The intracellular parasites (further cells. "schizonts", although also trophozoites were present) were subsequently released from the host cells saponin lysis (15 min in 0.1% saponin in CMF-Hanks at roomtemperature) and mechanical shearing.

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The schizonts were pelleted and separated from host material after centrifugation through 45% Percoll (Pharmacia Fine Chemicals) (20 min, 700g, 4°C). Dry pellets of schizonts were stored at -70°C until further use.

Triton X114 extraction

Triton X114 extractions were carried out to obtain the hydrophilic protein fraction of schizonts. The procedure used was described earlier by C. Bordier (1981) Journal of Biological Chemistry, vol. 256 no. 4 (feb) pp. 1604-1607.

10⁸ to 10⁹ E acervulina schizonts per ml of TBS (10 mM Tris-HCl, 150 mM NaCl pH7.4) were sonified ±3 x 20 sec. on ice with the microtip (Branson sonifier, position 7). PMSF (final concentration 1 mM) and DNase/RNase (final concentration for both 0.02 mg/ml) was added (DNase/RNase stock: 2 mg/ml DNase, 2 mg/ml RNase in 5 mM MgCl2).

Precondensed TritonX114 was added to the sonified schizonts in suspension to a final concentration of 10% (v/v) and mixed well to dissolve the proteins. The pelleted by non-extractable material was centrifugation 20 min 12,000g at 4°C. The soluble sucrose а cushion layered over fraction was sucrose, 0.06% (v/v) TX114 in TBS), incubated 10 min room temperature.

40°C and spun 10 min 400g at roomtemperature. The A 40°C and spun 10 min 400g at hydrophilic fraction was extracted again by the same procedure.

The hydrophilic fractions were stored at -70°C until further use. Total protein concentration was determined using the BCA (Pierce Chemicals) assay.

Prep-cell fractionation

Hydrophilic proteins were further separated with respect to their relative molecular mass on SDS-PAGE Laemmli the in reducing conditions under of preparative We made use Hereto electrophoresis in the so-called Prepcell.

Materials:

Propell Prep cell apparatus (Biorad Labs) with Prep cell

column (37mm ID) Drivs Dialyse membrane yse membrane for Prep cell (cut off 6kD)

A Power supply (EPS 600 Pharmacia)

Reducing sample buffer: 62.5 mM Tris-HCl pH 6.8; 10%

- A glycerol; 2% SDS; 0.01% bromophenol blue (Merck);
- 0.13 M DTT (dithiothreitol, Merck)

Electrophoresis buffer/elution buffer: 25 mM Tris, 192 mM Glycine, 0.1% SDS pH8.6

Method and results:

were performed at 4°C. For the procedures hydrophilic proteins fractionation of the stacking/9% separating gel (polyacrylamide) in the 37mm tube (filled to 6cm) of the Prepcell according to the manufacturers protocol, but with the addition of 0.1% SDS.

The hydrophilic phase of TX114 extractions kept at -70°C was thawed and the hydrophilic proteins (about 8mg per run) were diluted in reducing sample buffer (total volume was ± 6 ml), boiled 3 min 100°C, and were loaded on the surface of the 4% stacking gel using a narrow tube affixed to a syringe.

The Prepcell was connected to the powersupply and electrophoresis was started at 40mA, 500V max.

The collection of fractions (fraction volume ±2.5-3ml; flow 0.6ml/min) started after about 6 hours, when the tracking dye eluted from the cell. Fractions were collected overnight (±100 fractions) in plastic 3.5ml tubes (Sarstedt).

Samples of the fractions were taken for analysis by botton botton botton botton botton botton blotting. Fractions were stored at -70°C.

This purification method resulted in fractions containing almost pure proteins as follows from analyses shown below.

Amino acid sequencing

Prepcell run

A Prepcellrun COC9314612 Selected fractions of band around Mr=37kan almost pure containing (designated as EASC2) were pooled, concentrated by acetone precipitation and run on a 12% PAAgel. The gel was shortly stained with a non-denaturing Coomassie Brilliant Blue staining protocol: staining: 20 min at ambient temperature in 0.2% CBB in 20% methanol/0.5% acetic acid. Destaining: 60 min in 30% methanol.

The staining 37kD band was cut out. Internal amino acid sequencing was performed on a selected HPLC-purified peptide of a trypsin digest of the EASC2, all performed by Eurosequence BV Groningen The Netherlands.

The amino-acid sequence of the tryptic peptide was GWIKQEEVDDIVQK (see SEQ.ID.No:2 amino acids 212-225).

This coding sequence for this peptide was also detected after DNA sequencing of the clone.

A

EXAMPLE 2

Preparation of monospecific antibodies in rabbits

Prevaccination sera of SPF rabbits were screened on Weskin western blotted E. acervulina antigens of different developmental stages and on a blot of E coli proteins.

'Negative' rabbits were selected for the raising of antibodies.

Preparel runs

Fractions of Prepcellruns containing EASC2 (37kD) were selected by SDS-PAGE, pooled and concentrated (± 3x) with an Amiconcell (YM10 filter) to 3.5ml.

The rabbit was twice immunized with concentrated antigen in GNE (8x 0.25ml i.c.; 1ml i.p.) with an weeks after the second weeks. Two interval of 4 immunisation the rabbit was bled and sera were tested Wiskern blots of Eimeria acervulina en tenella sporozoites and schizonts 42hr. Figure 1 shows the immunodetection of the monospecific result of the antiserum on sporozoite antigens of both species. It appeared that the antibodies recognised a parasite product of about 37kD in both E. acervulina (Lane A1) and E. tenella (Lane B1). Control sera of the same rabbit prior to immunization did not recognise these bands (Lanes A/B2). The protein is also present in schizont stages of the two species (not shown).

EXAMPLE 3

<u>Vaccination of chickens with E.acervulina TX114</u> hydrophilic fraction and EASC2

The TX114 hydrophilic phase of schizont material was separated and dialysed extensively against 0.01M PBS pH 7.3 at 4°C.

Selected fractions containing the EASC2 37kD protein were dialysed extensively against 3 \times 5 liter 0.01 M PBS pH 7.3 at 4°C.

concentration of protein in the vaccine The by staining different estimated preparations was concentrations of sample with CBB after SDS-PAGE and with intensity of the staining comparing the reference sample of BSA.

The volumes were corrected to obtain \pm 5 μ g protein/dose for the purified protein and about 15 μ g/dose for the total hydrophilic fraction.

These were stored as aliquotted volumes for priming and booster vaccination at -70°C. Frozen vaccine preparations were thawed.

To every 15 ml of vaccine 3.2 mg Quil A Superfos Biosector was added as adjuvant in a volume of 1 ml 0.01 M PBS pH 7.3.

Vaccine was mixed well by vortexing and injected in 4-6 week old coccidia-free White Leghorn chickens in 0.75 ml given subcutaneously.

The vaccine contained 150 μ g Quil A/dose.

Figure 2 shows a Coomassie BB stained SDS-PAGE of the EASC2 (Lane 1) and 42hr TX114 hydrophilic fraction (Lane 2) injected into the chickens as vaccine.

Four weeks after priming birds were boosted with booster vaccine.

A the same dose via the same route. The booster-vaccine was prepared freshly from the frozen antigen stock.

Control chickens were inoculated with 150 μ g Quil A/dose in PBS. Each group comprised 14 chickens.

Eleven days after the boosting vaccination all chickens were inoculated orally with 240 sporulated oocysts of Eimeria acervulina H in 1 ml of 15% sucrose in water.

Chickens were placed in cages 2 birds per cage. Occyst output was assessed in fecal samples taken from days 4 to 8 after challenge.

Table 2 shows the results of this experiment. Occyst output is expressed as % occysts from the output in the control animals.

Statistical evaluation of the data was performed on the LOG of the number of oocysts using Student's Ttest or Mann-Whitney 's test if data distribution was not normal.

When p<0.05 the difference was regarded significant.

This table shows that both the TX114 fraction and the EASC2 preparell purified fraction induce a statistically significant reduction (p<0.05) in occyst output after challenge.

Prepcell

Prep cell purification seemed to improve the protection induced by the TX114 vaccine.

Table 2. Oocyst output in percents from control and statistical value of difference

Immunogen	% oocyst output	p value different			
	from control ± S.D.	from control			
EASC2 prep cell	72 ± 30	p=0.01			
pure ± 5 μg/dose		,			
Hydrophilic TX114	84 ± 17	p=0.02			
proteins of					
Schizonts ± 15					
μg/dose					

In another experiment in which only total extracts of 42 hr schizonts were used as vaccine no significant occyst reduction could be induced (results not shown).

In a second experiment prepectl purified EASC2 was used in two different concentrations of 0.2 and 2 µg/dose. Following the same protocol for immunization and challenge, protection was measured in ten chickens per group as reduction of oocyst output compared to the group inoculated with PBS/QuilA.

Table 3 summarises the average percentual oocyst output of the control for the two EASC2 vaccinated EASC2 groups. This table demonstrates that the dose dependent manner showing protected in a statistically significant difference at a dose of 2 μg/dose.

Table 3. Occyst output in percents from control and statistical value of difference

	% oocysts ± S.D.	significance of difference
Group	(control	from control(p-value)
	output=100%)	
EASC2/Quil A	64.0 ± 22	0.008
2 μg/dose		
EASC2/Quil A	90.2 ± 27	NOT SIGNIFICANT
0.2 μg/dose		

A

EXAMPLE 4

Immunological stimulation after vaccination with EASC2 or TX114 hydrophilic proteins.

In both protection experiments mentioned above chickens were assayed for stimulation of immunological parameters such as T-lymphocyte proliferation and serum antibodies.

Serum antibodies

Antibodies recognising the vaccine constituents were only detected in sera from the groups vaccinated with the 42hr TX114-hydrophilic fraction and not the group vaccinated with the purified EASC2.

Lymphocyte proliferation

Lymphocyte proliferation after antigenic stimulus was tested in a lymphocyte stimulation test (LST).

Method:

blood cells

Prior to challenge peripheral bloodcells were taken from all chickens of each group.

Peripheral blood leucocytes (PBL) were isolated by centrifugation 3 ml of the total blood for 7 min at ambient temperature. The buffy coat collected in RPMI 1640 (Dutch modification) and washed two times. Cell concentration was adjusted to 1x107 The RPMI 1640 (Dutch cells per ml in RPMI 1640. with sodium used was supplemented modification) pyruvate (1 mM), Glutamine (2 mM), penicillin 200 U/ml and streptomycin 200 μ g/ml.

96 well round-bottom tissue culture plates were seeded with 0.05 ml cell suspension with 3.0% chicken serum (Gibco BRL), 0.05 ml "stimulating antigen" suspension and 0.05 ml RPMI 1640, cultured for 64 hr at 41°C under 5% CO₂ atmosphere. Subsequently 18.5 kBq

tel-

3-H-Thymidine (Amersham Beckenham U.K.) was added per well and 8 hrs later the cells were harvested on a glass-fibre filter (Skatron Norway Bluemat) using a 96 well Cell Harvester (Skatron Norway). The filters were saturated with scintillation fluid (LKB BetaScint) and counted in a Betaplate 1205 (Pharmacia / LKB Sweden).

As "stimulating antigen" E.acervulina schizonts were used, which were sonicated using a microtip-equipped Branson sonifier at position 6 for 3x20 Stored Stored at -70°C. The antigens were thawed before use and diluted to meet the concentration used for the stimulation. PBL of all groups were stimulated with 3.10⁵ E. acervulina schizonts.

Statistical evaluation was performed using Student's T-test on the LOG of the Stimulation Index (SI) (the number of counts per min (cpm) of the stimulated cultures divided by the cpm of the non-stimulated control). When p<0.05 the difference was regarded significant.

Results:

Table 4. shows the mean S.I. for the groups from both experiments described above. The first experiment in which EASC2 vaccine was compared with the TX114-hydrophilic fraction, and the second experiment dealing with the two dosages of the EASC2 vaccine.

It appeared that all antigens or dosages induced a significant positive T-cell response detectable in the peripheral blood at the time of challenge.

In both experiments, however, the higher dose of the prepedl pure EASC2 vaccine (2 or 5 μ g/dose) induced the very highest stimulation of T-cells. The ranking of the T-cell stimulation correlated with the reduction in oocyst output after challenge.

Table 4. Mean incorporation of 3H -Thymidine after stimulation with E. acervulina schizonts by PBL from groups immunised with the different vaccines, expressed as Stimulation Index (S.I) \pm Standard Error (SE).

Experiment	Group	³ H-thymidine					
1		incorporation in					
	·	S.I. ± SE					
I	EASC2 5 μg	120 ± 47 @					
	TX114 hydrophilic proteins	31 ± 12 @					
	Placebo	6 ± 1					
II	EASC2 2µg	112 ± 28 @					
	EASC2 0.2 μg	24 ± 4 @					
	Placebo	2.3 ± 0.3					

@) Significant from controle group p<0.001

EXAMPLE 5 CLONING EXPERIMENTS

Sporulation of E. acervulina oocysts

A suspension of 5*10⁸ E. acervulina oocysts in 60 ml 10⁻⁴M sodium dithionite was centrifuged, after which the pellet was washed once with 100 ml sterile water. The cells were resuspended in 500 ml 2% potassium bichromate and then incubated under the influence of strong aeration for 7 hours at 30°C. The oocysts were then collected by centrifuging and washed three times with 200 ml sterile water.

Isolation of RNA

For the isolation of RNA (Pasternak J. et al., Mol. & Bioch. Par. 3, 133-142, 1981) the cell pellet was taken up into 2.8 ml of buffer containing 10 mM Tris acetate (pH7.6), 75 mM sodium acetate, 1% SDS, 2 mM EDTA, 0.2 mg/ml proteinase K and 10 mM vanadyl ribonucleoside complexes. The oocysts were destroyed by vortexing for 60 seconds (max) in the presence of 13 g glass beads (ø 0.5mm). 5 ml of phenol was added to the total extract and the mixture was vortexed for a centrifuging, the 60 seconds. After further pipetted off and again supernatant liquor was volume of with equal extracted an phenol/chloroform/isoamyl alcohol (25:24:1). RNA was precipitated after adding 2.5 volume ethanol and the resulting precipitate was dissolved in 800 ml Tris 10 mM, EDTA 0.1 mM pH 7.6 $(T_{10}E_{0.1})$, after which the product was extracted a further twice with an equal volume of phenol/chloroform/isoamyl alcohol (25:24:1) and twice with chloroform/isoamyl alcohol (24:1) and PolyA+-RNA ethanol. was with then precipitated of oligo(dT)-cellulose isolated by means chromatography (Maniatis T. et al.: Molecular Cloning. Cold Spring Harbor Laboratory, 1982). Approximately 100 μ g polyA⁺-RNA was isolated from 5*10⁸ oocysts.

cDNA synthesis

PolyA+-RNA was converted to cDNA by means of the enzyme MMLV reverse transcriptase. For this purpose 25 μ g polyA⁺-RNA was dissolved in 90 ml of water and denatured for 5 minutes at 20°C by adding mercury which Bafter 10 mM, methyl hydroxide to Mercaptoethanol was added to 45 mM and the mixture incubated for a further 3 minutes at 20°C. The enzyme reaction was carried out in 190 ml buffer containing 4 mg oligo(dT)15, 150 U RNasin(R), 20 mM Tris (pH 7.6), 30 mM KCl, 4 mM dithiothreitol (DTT), 2 mM MgCl2, 1 mM of each dNTP and 3000 U MMLV reverse transcriptase.

The reaction was stopped after 1 hour incubation at 37°C by adding 10 ml 0.5 M EDTA. After extraction with an equal volume of phenol/chloroform/ isoamyl alcohol the RNA/DNA hybrid was precipitated by (25:24:1), acetate to 2 M and 2.5 adding ammonium The combined action of the enzymes DNApolymerase I and RNase H (Gubbler U. et al., Gene 25, 263-269, 1983) results in the synthesis of the second string. The pellet was dissolved in 960 μ l of buffer containing 20 mM Tris (pH 7.6), 5 mM MgCl2, 100 mM 16 U RNase H, 200 U DNA- $(NH_4)_2SO_4$, 0.6 mM B-NAD, 20 U DNA-ligase (E.coli). polymeraseI and incubation time was 1 hour at 12°C and then 1 hour at 22°C, after which the reaction was stopped by adding an equal volume of phenol/chloroform/isoamyl alcohol (25:24:1) and precipitating with ethanol.

Before the cDNA was cloned in a vector suitable for this purpose it was first modified. cDNA (5 μ g) was dissolved in 100 μ l of buffer containing 30 mM sodium acetate (pH 5.6), 50 mM NaCl, 1 mM ZnSO4 and 21 U Mung Bean Nuclease. After incubation for 30 minutes at 37°C the reaction was stopped by adding EDTA to 10 mM and After extraction mM. to 25 phenol/chloroform/isoamylalcohol (25:24:1) the mixture Sephadex G50 desalinated over a column. following were added to the eluate (125 μ l): Tris pH 7.6 to 50 mM, EDTA to 2.5 mM, DTT to 5 mM, U adenosylmethionine to 0.5 mM, and 100 methylase. After incubation for 30 minutes at 37°C, the reaction was stopped by heating for 15 minutes at 65°C, after which 1/10 volume of a solution containing Tris-HCl 100 mM, MgCl₂ 100 mM and NaCl 500 mM (pH7.5) was added, and, at the same time, each dNTP to 1 mM and 12.5 U Klenow DNA-polymerase. The reaction was equal volume of stopped by adding an after alcohol (25:24:1) phenol/chloroform/isoamyl incubating for 60 minutes at 22°C. The supernatant

liquor was precipitated after adding 350 μ l H₂O and 50 with sodium acetate Hq) 5.6) $\mu 1$ isopropanol. After dissolving in 100 ml H₂O, the desalinated over Sephadex G50 and pellet was eluate precipitated with ethanol. After dissolving the pellet in 24 μ l H₂O, ligation was carried out in 50 μ l by adding 2 μ g EcoRI linker, Tris-HCl(pH 8.0) to 30 mM, MgCl₂ to 10 mM, dithiothreitol to 10 mM, ATP to 1 mM, qelatin to 0.1 mg/ml and 10 U T4DNA-ligase. reaction was stopped after 16 hours incubation at 4°C by heating (for 15 minutes at 70°C) after which cutting was carried out with restriction endonuclease EcoRI in 210 μ l buffer containing 100 mM Tris-HCl (pH 7.6), 50 mM NaCl, 10 mM MgCl₂, 2.5 mM DTT and 500 U minutes incubation After 90 at 37°C, reaction was stopped by means of extraction with an volume of phenol/chloroform/isoamylalcohol equal supernatant liquor was precipitated (25:24:1). The with 2.5 volume ethanol after adding sodium acetate (pH 5.6) to 300 mM cDNA and linkers were separated by of а Biogel A15M column. The cDNA was means precipitated with ethanol, after which the precipitate was dissolved in Tris-HCl 10 mM, EDTA 0.1 mM(pH 7.6). The cDNA molecules were then cloned in phage lambda ZAPII (Stratagene).

(2*10⁵ bank pfu) CDNA of the Screening antibodies directed against the EASC2 protein fraction of E acervulina schizonts revealed six positive phage clones. These antibodies were deluted 1:2000 with 1x Tris salt (Tris-HCl 10 mM, NaCl 150 mM, pH 8.0) + 0.05% Tween 20 + 10% Foetal Calf Serum (FCS) and incubated for two hours at room temperature (RT) with the filters. The filters were then washed 4 times, for 10 minutes each time, with 50 ml 1 x Tris salt + 0.05% filter. For the second antibody 20, each Tween incubation a conjugate of goat-anti-rabbit antibodies and alkaline phosphatase was used (diluted 1:7500 in

1x Tris salt + 0.05% Tween 20 + 10% FCS) and incubated for 30 minutes at RT, after which the filters were described after the first washed as incubation. The colour reaction was carried out in Tris-HCl 100 mM, NaCl 100 mM, MgCl₂ 10 mM, (pH 9.6), in which 0.33 mg/ml Nitrobluetetrazolium 5-bromo-4-chloro-3-indolyl phosphate dissolved. The filters were evaluated after 30 minutes incubation at RT. The immunopositive clones plaque-purified and rescued by means of in vivo according to the protocol of the excision, manufacturer (Stratagene). Plasmid DNA was isolated, the resulting in vivo excision clones, sequencing purposes according to standard protocols т., al. supra). Partial (Maniatis et sequence information showed all clones to be homologous, from largest clone the nucleotide sequence determined completely. This clone, designated pBLUE EASC2, contained an insert of 1566 bp.

Legend to the figures.

Fig. 1. Western blot of E. acervulina (A) and E. tenella (B) sporozoite proteins probed with antiserum raised against Prep cell purified EASC2 protein (Lane 1) or pre-immune control serum (Lane 2). Markers indicate molecular weight calibration in kD.

Fig. 2. Coomassie Brilliant Blue stained SDS-PAGE of Prep cell purified EASC2 protein (Lane 1) or TX114 hydrophilic fraction of E. acervulina 42hr schizonts (lane 2). Lane M contains molecular weight calibration markers in kD.

SEQUENCE LISTING

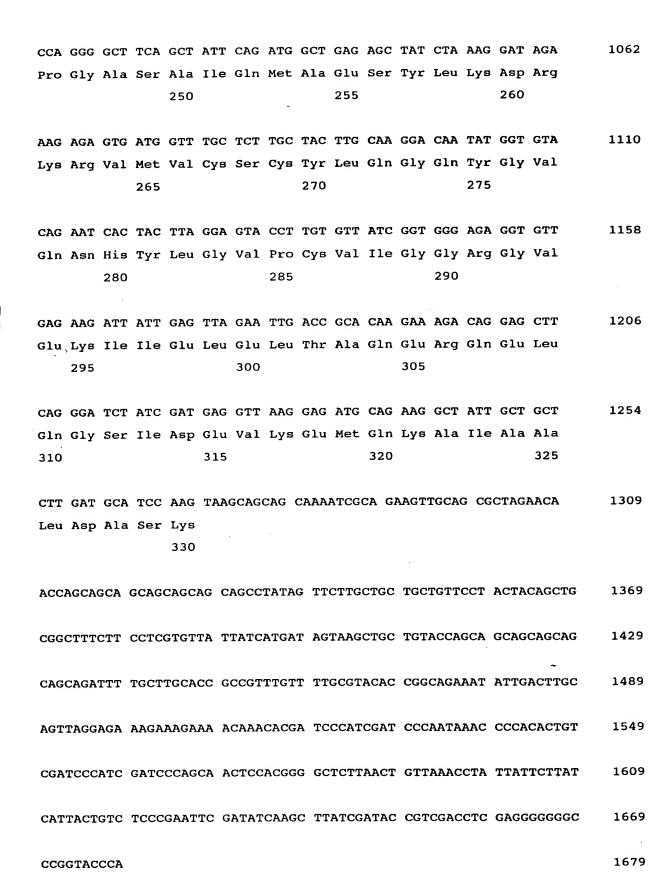
- (1) GENERAL INFORMATION:
 - (i) APPLICANT:
 - (A) NAME: Akzo Nobel N.V.
 - (B) STREET: Velperweg 76
 - (C) CITY: Arnhem
 - (E) COUNTRY: The Netherlands
 - (F) POSTAL CODE (ZIP): 6824 BM
 - (G) TELEPHONE: 04120-66204
 - (H) TELEFAX: 04120-50592
 - (I) TELEX: 37503 akpha nl
 - (ii) TITLE OF INVENTION: T cell stimulatory protein of Eimeria
 - (iii) NUMBER OF SEQUENCES: 2
 - (iv) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)
- (2) INFORMATION FOR SEQ ID NO: 1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1679 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA to mRNA
 - (iii) HYPOTHETICAL: NO

- (iv) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Eimeria acervulina
 - (D) DEVELOPMENTAL STAGE: Schizont
- (vii) IMMEDIATE SOURCE:
 - (B) CLONE: EASC2_1
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 280..1269
 - (D) OTHER INFORMATION:/function= "Eimeria lactate dehydrogenase"
- (ix) FEATURE:
 - (A) NAME/KEY: misc_feature
 - (B) LOCATION: 1..51
 - (D) OTHER INFORMATION:/label= pBluescriptII
- (ix) FEATURE:
 - (A) NAME/KEY: misc_feature
 - (B) LOCATION: 1624..1679
 - (D) OTHER INFORMATION:/label= pBluescriptII
- (ix) FEATURE:
 - (A) NAME/KEY: misc_feature
 - (B) LOCATION: 45..54
 - (D) OTHER INFORMATION:/label= EcoRI-linker
- (ix) FEATURE:
 - (A) NAME/KEY: misc_feature
 - (B) LOCATION: 1621..1630
 - (D) OTHER INFORMATION:/label= EcoRI-linker

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

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AAAC	TCTC	TA 1	TTTCC	CTCAT	IT AT	CTAC	CCGCI	TCF	ATCGC	STGG	GTGT	GTA	AGA (CGTAC	CGTACG	180
TACA	GCTC	GG (GCTGG	CTT#	C TO	CGC#	ACCGC	TT#	ATTT?	ATTA	CTT	ATTO	CAT A	ACACA	ATTTTA	240
TATO	TTTC	CTT (CTTCI	тттт	т ст	TGCI	CTTI	CTI	rgtg <i>i</i>					rrc (294
											1				5	
AAG	AAT	ACA	CGC	CCC	AAG	ATT	GCT	ATG	GTG	GGC	TCC	GGT	ATG	ATT	GGA	342
Lys	Asn	Thr	Arg	Pro	Lys	Ile	Ala	Met	Val	Gly	Ser	Gly	Met	Ile	Gly	
				10					15					20		
GGC	ACC	ATG	GCT	TTC	CTG	TGC	AGC	TTG	AGG	GAA	CTC	GGA	GAT	GTT	GTC	390
Gly	Thr	Met	Ala	Phe	Leu	Cys	Ser	Leu	Arg	Glu	Leu	Gly	Asp	Val	Val	
			25					30					35			
CTC	TTC	GAC	GTT	GTA	CCG	AAC	ATG	CCG	ATG	GGG	AAG	GCG	ATG	GAT	ATA	438
			Val													
		40					45			-	_	50				
TCG	CAC	AAT	TCG	TCG	GTG	GTT	GAC	ACG	GGT	ATA	ACA	GTA	TAC	GGC	TCA	486
Ser	His	Asn	Ser	Ser	Val	Val	Asp	Thr	Gly	Ile	Thr	Val	Tyr	Gly	Ser	
	55					60					65					
አአጥ	ጥሮል	ጥልሮ	CAC	TGC	ጥጥር	AAG	CCT	GCG	GAC	GTA	GTA	ATA	ATA	ACA	GCA	534
			Glu													
	ser	IYL	GIU	Cys	75	Буз	GIY	7124	no _P	80					85	
70					73										•	
GGG	АТА	ACA	AAG	АТА	CCC	GGA	AAG	AGC	GAT	AAA	GAA	TGG	TCT	AGA	ATG	582
			Lys													
				90					95					100		

GAT	CTA	TTA	CCT	GTG	AAT	ATA	AAG	ATA	ATG	AGG	GAG	GTC	GGT	GCA	GCA	630
Asp	Leu	Leu	Pro	Val	Asn	Ile	ГЛа	Ile	Met	Arg	Glu	Val	Gly	Ala	Ala	
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Ile	Lys	Ser	Tyr	Сув	Pro	Asn	Ala	Phe	Val	Ile	Asn	Ile	Thr	Asn	Pro	
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TTA	GAT	GTG	ATG	GTA	GCT	GCT	CTT	CAA	GAG	TCA	TCA	GGA	CTA	CCT	CAT	726
Leu	Asp	Val	Met	Val	Ala	Ala	Leu	Gln	Glu	Ser	Ser	Gly	Leu	Pro	His	
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					•											
CAT	AGA	ATC	TGC	GGT	ATG	GCT	GGG	ATG	CTT	GAT	AGC	TCT	CGT	TTT	AGA	774
His	Arg	Ile	Сув	Gly	Met	Ala	Gly	Met	Leu	Asp	Ser	Ser	Arg	Phe	Arg	
150					155					160					165	
													GTA			822
Arg	Met	Ile	Ala	Asp	Lys	Leu	Glu	Val	Ser	Pro	Arg	Asp	Val	Gln	Gly	
				170					175					180		
																ън.
													AGT			870
Met	Val	Ile	Gly	Val	His	Gly	Asp	His	Met	Val	Pro	Leu	Ser	Arg	Tyr	
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																242
													AAG			918
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																266
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Ile	Lys	Gln	Glu	Glu	Val	Asp	Asp	Ile	Val	Gln		Thr	Lys	Val	Ala	
	215					220					225					
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(2) INFORMATION FOR SEQ ID NO: 2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 330 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:
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- Lys Ala Met Asp Ile Ser His Asn Ser Ser Val Val Asp Thr Gly Ile
 50 55 60
- Thr Val Tyr Gly Ser Asn Ser Tyr Glu Cys Leu Lys Gly Ala Asp Val
 65 70 75 80
- Val Ile Ile Thr Ala Gly Ile Thr Lys Ile Pro Gly Lys Ser Asp Lys
 85 90 95
- Glu Trp Ser Arg Met Asp Leu Leu Pro Val Asn Ile Lys Ile Met Arg 100 105 110
- Glu Val Gly Ala Ala Ile Lys Ser Tyr Cys Pro Asn Ala Phe Val Ile 115 120 125

- Asn Ile Thr Asn Pro Leu Asp Val Met Val Ala Ala Leu Gln Glu Ser 130 135 140
- Ser Gly Leu Pro His His Arg Ile Cys Gly Met Ala Gly Met Leu Asp 145 150 155 160
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- Arg Asp Val Gln Gly Met Val Ile Gly Val His Gly Asp His Met Val
- Pro Leu Ser Arg Tyr Ala Thr Val Asn Gly Ile Pro Leu Ser Glu Phe
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- Val Lys Lys Gly Trp Ile Lys Gln Glu Glu Val Asp Asp Ile Val Gln 210 215 220
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- Gly Gly Arg Gly Val Glu Lys Ile Ile Glu Leu Glu Leu Thr Ala Gln 290 295 300
- Glu Arg Gln Glu Leu Gln Gly Ser Ile Asp Glu Val Lys Glu Met Gln 305 310 315 320

Lys Ala Ile Ala Ala Leu Asp Ala Ser Lys 325 330